## Schreiber, David

From:

Sent:

Steadman, David (AU1652) Monday, March 10, 2003 4:23 PM

To:

Schreiber, David

Subject:

- 09/991,212 sequence comparison

NAME: David Steadman

AU: 1652

Date:03/10/03 Office: 10D-04 Mailbox: 10D-01

Mr. Schreiber, please compare the following sequences:

1) SEQ ID NO:1 against SEQ ID NO:5

2) SEQ ID NO:2 against SEQ ID NO:5

" Please save results to diskette.

Thank you very much.

David J. Steadman Art Unit 1652 Crystal Mall 1 Room 10D-04 703-308-3934

Access DB	u
ALLESS DD	#

## **SEARCH REQUEST FORM**

## Scientific and Technical Information Center

Requester's Full Name:		Evaminar # .	Data
Art Unit:	Phone Number 30	Examiner # :Serial Number:	Date.
Mail Box and Bldg/Room I	ocation:	Results Format Preferred (circle):	PAPER DISK E-MAII
If mor than one search i	s submitted, please p	prioritize searches in order of ne	ed.
Please provide a detailed stateme Include the elected species or stre	ent of the search topic, and outures, keywords, synonymy terms that may have a s	describe as specifically as possible the sub ms, acronyms, and registry numbers, and c pecial meaning. Give examples or relevan	ject matter to be searched.
Title of Invention:			
Inventors (please provide full r	names):		
Earliest Priority Filing Date	::		
•		ormation (parent, child, divisional, or issued p	atent numbers) along with the
· .			
*****	******	*************	******
STAFF USE ONLY	Type of Search	Vendors and cost wh	ere applicable
Searcher: DSchvei		2_ STN	
Searcher Phone #: 308 -47		Dialog	
Searcher Location: CMI C	AU3 Structure (#)	. Questel/Orbit	**
Date Searcher Picked Up:	Bibliographic	Dr.Link	
Date Completed: 3/11	Litigation	Lexis/Nexis	
Searcher Prep & Review Time:	Fulltext	Sequence Systems Comp	ygen GCG
Clerical Prep Time:	Patent Family	WWW/Internet	
Online Time: 26	Other	Other (specify)	

PTO-1590 (8-01)

```
RESULT
HSZ83953
                              570 bp
                                      mRNA
                                             linear
                                                    PRI 16-JAN-1997
LOCUS
          HSZ83953
DEFINITION
         H.sapiens mRNA; clone CD 274.
ACCESSION
          283953
VERSION
          Z83953.1 GI:1783342
KEYWORDS
SOURCE
          human.
  ORGANISM
         Homo sapiens
          Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
          Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
REFERENCE
          1 (bases 1 to 570)
         Gasparini, P.
  AUTHORS
  JOURNAL
         Unpublished
REFERENCE
          2 (bases 1 to 570)
 AUTHORS
         Gasparini, P.
  TITLE
          Direct Submission
         Submitted (13-JAN-1997) Gasparini P., Servizio di Genetica Medica,
  JOURNAL
          I.R.C.C.S. - Ospedale CSS Viale Cappuccini, I-71013 San Giovanni
          Rotondo, Foggia, ITALY
FEATURES
                 Location/Qualifiers
                 1. .570
    source
                 /organism="Homo sapiens"
                 /db xref="taxon:9606"
                 /chromosome="6"
                 /map="6p22"
                 /clone="CD 274"
BASE COUNT
             127 a
                    134 c
                           134 g
                                   171 t
                                            4 others
ORIGIN
 Query Match
                     18.0%; Score 296; DB 9; Length 570;
 Best Local Similarity 100.0%; Pred. No. 1.2e-155;
 Matches 296; Conservative
                           0; Mismatches
                                         0; Indels
Qу
     519 GTGGGGCCCTCCACAAGAACGAAGCAGACTCTGCAGCATTGCTTTATCAGGAATGTTACT 578
        Db
     275 GTGGGGCCTCCACAGAACGAAGCAGACTCTGCAGCATTGCTTTATCAGGAATGTTACT 334
Qу
     579 GGGATGCTTTACTGCCATCCTCATAGGTGGCTTCATTAGTGAAACCCTTGGGTGGCCCTT 638
        335 GGGATGCTTTACTGCCATCCTCATAGGTGGCTTCATTAGTGAAACCCTTGGGTGGCCCTT 394
     Qу
        Db
        TGTCTTCTATATCTTTGGAGGTGTTGGCTGTCTCTCTCTGGTTTGTTGTGAT 454
Qу
     699 TTATGATGACCCCGTTTCCTATCCATGGATAAGCACCTCAGAAAAAGAATACATCATATC 758
        Db
     455 TTATGATGACCCCGTTTCCTATCCATGGATAAGCACCTCAGAAAAAGAATACATCATATC 514
     759 CTCCTTGAAACAACAGGTCGGGTCTTCTAAGCAGCCTCTTCCCATCAAAGCTATGC 814
Qу
        Db
     515 CTCCTTGAAACAACAGGTCGGGTCTTCTAAGCAGCCTCTTCCCATCAAAGCTATGC 570
```

```
seq documentation block:
          H$Z83953
LOCUS
                                570 bp
                                        mRNA
                                               linear PRI 16-JAN-1997
DEFINITION H. sapiens mRNA; clone CD 274.
ACCESSION
          Z83953
VERSION
          Z83953.1 GI:1783342
KEYWORDS
SOURCE
          human.
 ORGANISM Homo sapiens
          Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
          Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
          1 (bases 1 to 570)
REFERENCE
          Gasparini, P.
 AUTHORS
 JOURNAL.
          Unpublished
REFERENCE
          2 (bases 1 to 570)
 AUTHORS
          Gasparini, P.
 TITLE
          Direct Submission
          Submitted (13-JAN-1997) Gasparini P., Servizio di Genetica Medica,
  JOURNAL
          I.R.C.C.S. - Ospedale CSS Viale Cappuccini, I-71013 San Giovanni
          Rotondo, Foggia, ITALY
FEATURES
                  Location/Qualifiers
    source
                  1. .570
                  /organism="Homo sapiens"
                  /db xref="taxon:9606"
                  /chromosome="6"
                  /map="6p22"
                  /clone="CD 274"
BASE COUNT
             127 a
                    134 c
                            134 g
                                    171 t
                                              4 others
ORIGIN
alignment_scores:
          Quality:
                   98.00
                                   Length:
                                             98
           Ratio: 1.000
                                    Gaps:
Percent Similarity: 100.000
                          Percent Identity: 100.000
alignment block:
US-09-991-212A-1 x HSZ83953
Align seg 1/1 to: HSZ83953 from: 1 to: 570
     96 TrpGlyProProGlnGluArgSerArgLeuCysSerIleAlaLeuSerGl 112
       1111121212
    276 TGGGGCCCTCCACAAGAACGAAGCAGACTCTGCAGCATTGCTTTATCAGG 325
    112 yMetLeuLeuGlyCysPheThrAlaIleLeuIleGlyGlyPheIleSerG 129
    129 luThrLeuGlyTrpProPheValPheTyrIlePheGlyGlyValGlyCys 145
       376 AAACCCTTGGGTGGCCCTTTGTCTTCTATATCTTTGGAGGTGTTGGCTGT 425
    146 ValCysCysLeuLeuTrpPheValValIleTyrAspAspProValSerTy 162
       426 GTCTGCTGCCTTCTCTGGTTTGTTGTGATTTATGATGACCCCGTTTCCTA 475
    162 rProTrpIleSerThrSerGluLysGluTyrIleIleSerSerLeuLysG 179
       476 TCCATGGATAAGCACCTCAGAAAAAGAATACATCATATCCTCCTTGAAAC 525
    179 lnGlnValGlySerSerLysGlnProLeuProIleLysAlaMet 193
       526 AACAGGTCGGGTCTTCTAAGCAGCCTCTTCCCATCAAAGCTATG 569
```

## -continued

(B) CLONE: XLR

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:6:

CTTGATGCTC CCATGAGAAA ACTGG

25

- (2) INFORMATION FOR SEQ ID NO:7:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 25 base pairs (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear
  - (vii) IMMEDIATE SOURCE:
    - (B) CLONE: XLF
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:7:

AGGATTTTCG AGCATAGCAC CTGTC

25

What is claimed is:

- 1. An isolated and purified polynucleotide fragment encoding a polypeptide comprising the amino acid sequence of SEQ ID No:1.
- 2. A hybridization probe comprising the polynucleotide fragment of claim 1.
- An isolated and purified polynucleotide fragment comprising SEQ ID No:2.
- 4. An isolated and purified polynucleotide fragment which 30 is completely complementary to the polynucleotide of claim 1.
- 5. A hybridization probe comprising the polynucleotide fragment of claim 4.
- 6. An expression vector comprising the polynucleotide of 35 claim 1.
  - 7. A host cell containing the expression vector of claim 6.
- 8. A method for producing a polypeptide comprising the amino acid sequence of SEQ ID NO:1, the method comprising the steps of:

- a) culturing the host cell of claim 7 under conditions suitable for the expression of the polypeptide; and
- b) recovering the polypeptide from the host cell culture.
- 9. A method for detection of a polynucleotide encoding a polypeptide comprising the amino acid sequence of SEQ ID NO:1 in a biological sample containing nucleic acid material, the method comprising the steps of:
  - a) hybridizing the polynucleotide of claim 4 to the nucleic acid material of the biological sample, thereby forming a hybridization complex; and
  - b) detecting the hybridization complex, wherein the presence of the hybridization complex correlates with the presence of the polynucleotide encoding the polypeptide in the biological sample.

\* \* \* \* \*